AD			

Grant NUMBER DAMD17-94-J-4386

TITLE: Detection and Characterization of Autoantigens

in Breast Cancer

PRINCIPAL INVESTIGATOR: Janis Racevskis, Ph.D.

CONTRACTING ORGANIZATION: Montgomery County, Maryland

Rockville, Maryland 20850

REPORT DATE: October 1996

TYPE OF REPORT: Annual

PREPARED FOR: Commander

U.S. Army Medical Research and Materiel Command Fort Detrick, Frederick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for public release;

distribution unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

19970326 013

REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Warden, to Wa

1. AGENCY USE ONLY (Leave blank	k) 2. REPORT DATE	3. REPORT TYPE AN					
	October 1996	Annual (15 Ju	l 95 - 14 Jul 96)				
4. TITLE AND SUBTITLE			5. FUNDING NUMBERS				
Detection and Characte							
Cancer	DAMD17-94-J-4386						
			1				
6. AUTHOR(S)							
Janis Racevskis, Ph.D	•						
7. PERFORMING ORGANIZATION N	IAME(S) AND ADDRESS(ES)		8. PERFORMING ORGANIZATION REPORT NUMBER				
Montgomery County, Mar	ard and		REPORT HOMBER				
Rockville, Maryland 2							
Nochville, haryrana 2	.0030						
9. SPONSORING/MONITORING AGE	ENCY NAME(S) AND ADDRESS(ES)		10. SPONSORING/MONITORING				
Commander			AGENCY REPORT NUMBER				
	earch and Materiel Comm						
Fort Detrick, Frederic	ck, Maryland 21702-501	L2	:				
·							
			1				
11. SUPPLEMENTARY NOTES							
12a. DISTRIBUTION / AVAILABILIT	Y STATEMENT		12b. DISTRIBUTION CODE				
TEL DIGITIES HOLY ATAISASIEN	· O.A.L.		125. DISTRIBUTION CODE				
Approved for public re	elease; distribution un	nlimited					
13. ABSTRACT (Maximum 200	_						
Characterization of	of two breast tumor	c associated	autoantigens, which				
are newly discover	red gene products,	has revealed	that the first one				
(Ngp-1) is a GTP-R	oinding protein whi	ch localizes	exclusively to the				
	nteracts with rib						
interactions with	other proteins ar	re being inv	estigated using the e is located on the				
short arm of huma	vector system. In	t position 1	p34-1p35. A clone				
containing the ful	1 length transcript	of the secon	nd autoantigen clone				
Auag2 has been o	obtained and is h	peina semien	ced to completion.				
Recombinant protei	n Auag2 has been pro	oduced and pu	rified for antiserum				
production. No p	art of the Auag2 s	sequence is	found in any of the				
databases, althou	logy to a vascular						
endothelial growth	factor. Additiona	l breast cand	er patient sera have				
been collected and will be tested for autoantibodies.							
14. SUBJECT TERMS	15. NUMBER OF PAGES						
Breast Cancer, Autoa	15						
			16. PRICE CODE				
17. SECURITY CLASSIFICATION	18. SECURITY CLASSIFICATION	19. SECURITY CLASSII	FICATION 20. LIMITATION OF ABSTRACT				
OF REPORT	OF THIS PAGE	OF ABSTRACT	Similar of Aborrace				
Unclassified	Unclassified	Unclassified	IInlimited				

FOREWORD

Opinions, interpretations, conclusions and recommendations are those of the author and are not necessarily endorsed by the US Army.

Where copyrighted material is quoted, permission has been obtained to use such material.

Where material from documents designated for limited distribution is quoted, permission has been obtained to use the material.

Of Citations of commercial organizations and trade names in this report do not constitute an official Department of Army endorsement or approval of the products or services of these organizations.

In conducting research using animals, the investigator(s) adhered to the "Guide for the Care and Use of Laboratory Animals," prepared by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Resources, National Research Council (NIH Publication No. 86-23, Revised 1985).

For the protection of human subjects, the investigator(s) adhered to policies of applicable Federal Law 45 CFR 46.

In conducting research utilizing recombinant DNA technology, the investigator(s) adhered to current guidelines promulgated by the National Institutes of Health.

In the conduct of research utilizing recombinant DNA, the investigator(s) adhered to the NIH Guidelines for Research Involving Recombinant DNA Molecules.

In the conduct of research involving hazardous organisms, the investigator(s) adhered to the CDC-NIH Guide for Biosafety in Microbiological and Biomedical Laboratories.

David Rachani 10/24/96
PI - Signature Date

RACEVSKIS, Janis

TABLE OF CONTENTS

	PAGE
FRONT COVER	1
REPORT DOCUMENTATION PAGE	2
FOREWORD	3
TABLE OF CONTENTS	4
INTRODUCTION	5
BODY	6 - 7
CONCLUSIONS	8
REFERENCES	9
APPENDICES	10-15

INTRODUCTION

Tumor growth is associated with the expression of mutated gene products, inappropriate gene expression, and the breakdown of tissue architecture, leading to the exposure and release into the peripheral circulation of sequestered antigens (1,2). Whether these circulating, mutated or newly displayed tumor-associated antigens elicit an autologous humoral immune response in the breast tumor patient is of vital interest. Isolation, identification and characterization of novel breast tumor associated autoantigens might yield new insights into the disease process, and moreover, may be developed into diagnostic screening tests and potential targets for immunotherapy.

The screening of cDNA expression libraries with autologous patient serum is a powerful technique, which has been used successfully for the identification of autoimmune disease antigens (3), and which we have adapted for the identification of autoantigens in cDNA libraries made from breast tumor mRNA. After screening cDNA libraries, derived from primary ductal breast carcinomas with autologous patient serum, we have detected and isolated two immunoreactive cDNA clones, both of which are newly discovered gene We propose to fully characterize the identified autoantigens and to construct additional cDNA libraries and screen them with autologous serum to identify and isolate additional breast tumor autoantiqen cDNAs. The ultimate goals of our research project are: 1. To isolate autoantique clones which individually or in combination react specifically with most breast tumor patient sera and may form the basis for the development of diagnostic tests or perhaps identify potential targets for immunotherapy, and, 2. To test the hypothesis that breast tumors result in the expression of a characteristic profile of autoantigens.

BODY

During the past year we have concentrated most of our research effort at characterizing the two autoantigenic gene products that we identified, both of which are newly discovered genes. The most progress has been made in the characterization of the first of our isolated breast tumor autoantigens, and the results of our initial studies have been published (4). In addition, a figure from our paper showing nucleolar staining with anti autoantigen antibody was chosen as the cover illustration for the february 1996 issue of Cell Growth and Differentiation (appendix, page 10).

We have named the gene encoding the first autoantigen Ngp-1, (Genbank accession # L05425), and from the predicted amino acid sequence, have determined that it encodes a GTP-binding protein (5,6). Immunohistochemical analysis of tissue sections with affinity purified antiserum raised against a recombinant Ngp-1 protein revealed that the antigen was exclusively localized in the nucleolus and nucleolar organizer regions in all cell types analyzed (hence our proposed name Ngp-1: Nucleolar G-Protein gene The arrangement and spacing of the GTP binding protein motifs indicate that Ngp-1 belongs to a newly described subfamily of GTPases with one other known human member (HSR1) (7), the others The alignment of the GTP-binding being of prokaryotic origin. protein motifs of Ngp-1 with those of other members of the new subfamily are shown in the appendix, page 11. The discoverers of (Dr. Pontarotti et al, Centre National de la Recherche Scientifique, Toulouse, France) contacted me to request a full length probe for Ngp-1, and suggested a collaboration to determine the chromosomal location of the Ngp-1 gene. The results indicated that the Ngp-1 gene is located on the short arm of human chromosome 1, in the 1p35-1p34 region (appendix, page 12).

Southern blot analysis of Eco R1 digested human genomic DNA and hybridization against a full length Ngp-1 cDNA probe revealed two major bands hybridizing (approximately 15 and 8 kb in length). Two fainter bands were detected at 6 and 2.5 kb as well (appendix, page 13). Subcloning these genomic restriction fragments into standard vectors would be straight forward, however, sequencing this length of DNA would be a major timeconsuming undertaking which we will not begin at this time, since we feel there are more important projects to complete.

Since all GTPases interact with other cellular macromolecules (6), we set out to identify other gene products which interact with Ngp-1 during its regulatory functions. To accomplish this we have subcloned the entire open reading frame portion of Ngp-1 into

phagemid vector pBD-GAL4 (the bait plasmid) which we have begun to test in the yeast two-hybrid vector system. The yeast two-hybrid vector system is one of the most efficient techniques available for detecting in-vivo protein interactions (8). In a request for anti Ngp-1 antibody from a researcher in Germany (Dr. Stephan Witte, University of Konstanz) I learned that Ngp-1 protein interacted strongly with human ribosomal protein L7. Human ribosomal protein L7 is itself an autoantigen, and is associated with Systemic Lupus Erythematosus (9). The interaction was observed by Dr. Witte using the yeast two hybrid vector system with human ribosomal protein L7 as the bait.

We are continuing our work on characterization of our second breast tumor autoantigen isolate (working name Auag2), which is also a newly discovered gene. No part of this gene is to be found in any of the gene sequence databases. This is surprising considering the recent proliferation of partial cDNA sequence data entered in the databases from entire cDNA libraries. A possible explanation for the absence of Auag2 sequence data in the databases is that Auag2 contains regions of extremely high GC content (appendix, page 14), probably making reverse transcription of the mRNA difficult because of secondary structure, hence Auag2 is under-represented in cDNA The high GC content also makes accurate sequencing a difficult task, we have however deposited a partial sequence of Auag2 in the Genbank database (accession # U24576). finally able to isolate a clone containing 2.1 kb of Auag2 sequence (approximately the size of the Auag2 mRNA as determined by northern blot) by using the Gene Trapper technology (Life Sciences), where a cDNA library in a plasmid vector is converted to single strand hybridized with gene specific biotinvlated then oligonucleotide probes which are captured by avidin coated magnetic beads, thus highly enriching for the desired gene product. northern blots of mRNAs isolated from various human tissues, Auag2 appears to be most highly expressed in testes and brain; and is not detectable in liver and kidney (appendix, page 15). In certain tissues such as skeletal muscle there appears to be an extra band; perhaps representing different splicing products. Recombinant Auag2 protein has been produced and has been purified for antiserum production, which will be used for immunohistochemical localization of Auag2 protein within different tissues and cell types.

In the last progress report we reported that we had isolated a third potential autoantigen, however this proved to be non-reproducible. In the interim we have collected additional serum samples from breast cancer patients, and plan to screen these for the presence of autoantibodies, reacting with specific autoantigens.

CONCLUSIONS

We have made considerable progress in characterizing autoantigen Ngp-1, which we have found to be a very unusual molecule because it is one of the few GTP-binding proteins with a nuclear localization. We have determined the chromosomal location of the Ngp-1 gene, and learned that it interacts with ribosomal protein L7, itself a known autoantigen. Judging by the number of requests for probes and antisera there is great interest in this gene product. The second autoantibody clone Auag2 has an unusually high GC content and is under-represented in cDNA libraries. We were able to isolate a full length clone using a hybridization affinity technique. Although it shows some homology to vascular endothelial growth factor in one region, it does not match up with anything presently in the databases.

REFERENCES

- 1. Naftzger, C., and Houghton, A.N. Tumor immunology. Current Opinion in Oncology, 3:93-99, 1991.
- 2. Henderson, R.A., and Finn, O.J. Human tumor antigens are ready to fly. Advances in Immunology, 62:217-251, 1996.
- 3. Tan, E.M. Autoantibodies in pathology and cell biology. Cell, 67:841-842, 1991.
- 4. Racevskis, J., Dill, A., Stockert, R., and Fineberg, S.A. Cloning of a novel nucleolar guanosine 5'-triphosphate binding protein autoantigen from a breast tumor. Cell Growth & Differentiation, 7:271-280, 1996.
- 5. Saraste, M., Sibbald, P.R., and Wittinghofer, A. The P-loop, a common motif in ATP- and GTP-binding proteins. Trends Biochem. Sci., 15:430-434, 1990.
- 6. Bourne, H.R., Sanders, D.A., and McCormick, F. The GTPase superfamily: conserved structure and molecular mechanism. Nature, 349:117-127, 1991.
- 7. Vernet, C., Ribouchon, M.-T., Chimini, G., and Pontarotti, P. Structure and evolution of a member of a new subfamily of GTP-binding proteins mapping to the human MHC class I region. Mammalian Genome, 5:100-105, 1994.
- 8. Fields, S., and Ok-kyu Song. A novel genetic system to detect protein-protein interactions. Nature (London), 340:245-247, 1989.
- 9. von Mikecz, A., Hemmerich, P., Peter, H.H., and Krawinkel, U. Characterization of eukaryotic protein L7 as a novel autoantigen which frequently elicits an immune response in patients suffering from systemic autoimmune diseases. Immunobiol, 192:137-154, 1994.



Cover illustration of February 1996 issue of Cell Growth and Differentiation, showing immunohistochemical stain of a human testis section, with antibody against Ngp-1 (4). In the cross section of the seminiferous tubule shown, prominent staining is present in the nucleoli of cells nearer the basement membrane (spermatogonia, primary spermatocytes and Sertolli cells) as well as in interstitial Leydig cells.

C. Vernet et al.: New GTP-binding protein on MHC class I

G4

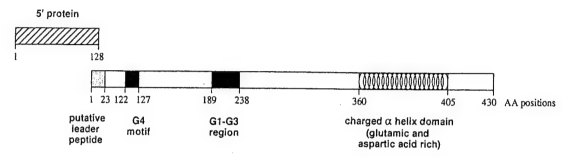


Fig. 1. Schematic representation of the whole protein HSR1. The 5' ORF protein, overlapping the seven first amino acids of the HSR1 protein, is also presented.

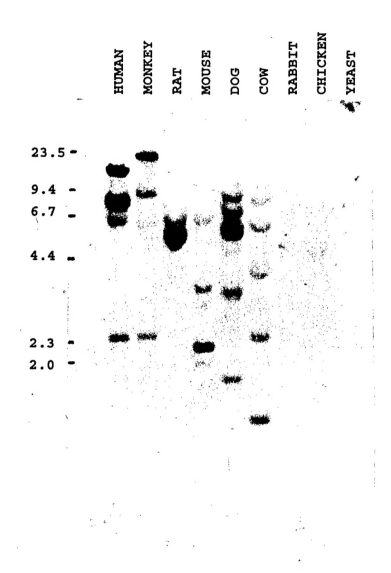
Ngp-1	IGYPNVGKSS VIN	D C P G	v
HSR1 (189-238) MMR1 (189-238) YRB1 HALCU (67-118) Obg BACSU (164-216) Bra ECOLI (14-76) THDF BACSU (227-279) THDF PSEPU (222-274)	I G F P N V G K S S L I N G V G F P S V G K S T L L S V V G P P S V G K S T L L N K I G R P N V G K S T L L N K I G R P N V G K S S L L N S	LLVGRKV VSVSRTPGHTR-YFQTYFLT PSVKLCDCPG LVGRKV VSVSRTPGHTR-YFQTYFLT PSVKLCDCPG MTNADS SVGAYEFTTL NVNPGMLEY- RGANIQLLDVPG VSSAKP KIADYHFTTL VPNLGMVETD DGRSFVMAD LPG LLGQKI SILTSRKAQTT RHRIVGIHTE GAYQAIYVDTPG LLVHEAK AIVTDIPGTT RDVIEEYVNV RGVPLRLVDTAG	LLLL
:Consensus	V G F P N V G K S S L L N .	L	T.
1	G1	G3	-
Ngp-1	FVLNKCDL		
HSR1 (45-52) MMR1 (45-52) YRB1 HALCU (240-247) Obg BACSU (339-346) Era ECOLI (121-128) THDF BACSU (333-339) THDF FSEPU (333-339)	L V L N K V D L L V L N K V D L V T V N K V D L I V A N K M D M L A V N K V D N V I L N K T D L L I R N K A D L		
Consensus	N K . D L	Fig. 2. Polypeptide sequence similarity between the different members of the new GTF binding protein family. The amino acid sequences (one-letter symbols) are aligned for maximal homology. Amino acids identical to the corresponding residues in the HSR I	

sequence are boxed.

Alignment of GTP-binding protein motifs of Ngp-1 with those of other sub-family members. Ngp-1 motifs are superimposed on figure from Vernet et al (7).



In-situ hybridization map of human chromosome 1, using a cDNA probe of Ngp-1, showing localization of signal to region 1p35-1p34.



Southern blot containing $4\mu g$ of EcoR I restriction digested genomic DNA per lane from nine eukaryotic species, hybridized against an Ngp-1 probe. Species DNA in each lane is identified on top of blot, and position of $\lambda/{\rm Hind~III}$ DNA size markers is noted on the left side of the blot.

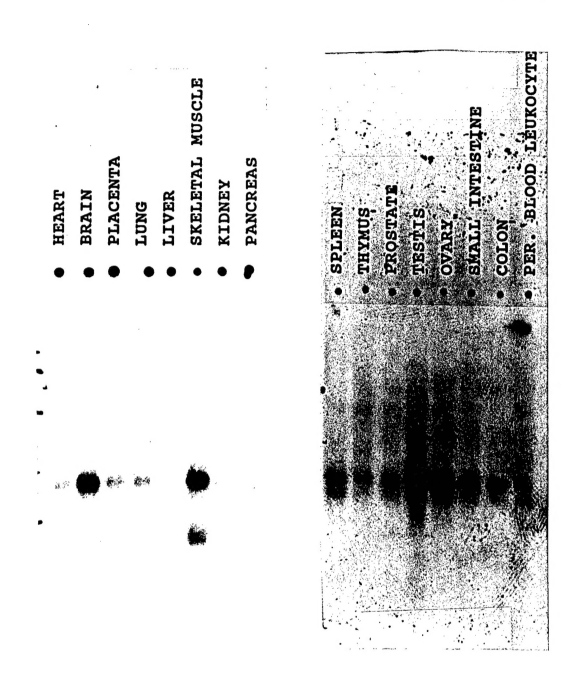
FILE NAME : AUTAG2.DNA

SEQUENCE: 587BP; 91 A; 214 C; 224 G; 58 T

*** SEQUENCE LIST *** (SINGLE)

TCGCGGAGGGAGCGAGCCGGCTAGAGGCCAGCTCCGCCGCCGCCGCCGCCTCCGAG CGCGGCCGCTGTTGTGTCTGCGACTGCTCCCGGCCGGAGGTGCAGGGAGCTCAGCCGAGC TCAAAGTGAAGCCACATTTGCCAAACTTGCAGCAGCGATTCGCAGCAGTTGCTGCCGCTG CGGCCGCGCCTGAAGCCGCGCGCGCGCGCGCGAGGGCTCCTGCAGCTGCGTCGCGCGC AGTCGGAGGCGGAGAAGGACGAAGACTGAGACTTCTGTCTCCCGGCCCCCCGG CACTTACGACGGGGGCCCCCAACCCGCCCCAGAGCAACGGCGATTT 3'

Sequence of original autoantigenic Auag2 isolate cDNA showing extremely high GC content (Genbank accession # U24576).



Northern blots containing approximately $2\mu g$ of poly A^+ RNA per lane from sixteen different human tissues (identified on top of blots) hybridized against an Auag2 probe. Location of RNA size marker bands is indicated in the left margin of the blot; approximate size of major band is 2.1 kb.